### **EXHIBIT RAS-3**

This is exhibit RAS-3 referred to in Declaration Under 37 C.F.R. 1.132 by Richard Anthony

Strugnell dated

Richard Strugnel

# Best Available Copy

## Use of peptide synthesis to probe viral antigens for epitopes to a resolution of a single amino acid

(antigenic determinant/foot-and-mouth disease virus)

H. MARIO GEYSEN\*, ROB H. MELOENT, AND SIMON J. BARTELINGT

92 \*Commonwealth Serum Laboratories, 45, Poplar Road, Parkville, Melbourne, Australia 3052; and †Central Veterinary Institute, 39 Houtribweg, Lelystad, The Netherlands raine and a subject to the contract of the con

Communicated by G. J. V. Nossal, March 12, 1984

ABSTRACT A procedure is described for rapid concurrent synthesis on solid supports of hundreds of peptides, of sufficient purity to react in an enzyme-linked immunosorbent assay Interaction of synthesized peptides with antibodies is then easily detected without removing them from the support. In this manner an immunogenic epitope of the immunologically important coat protein of foot-and-mouth disease virus (type Oi) is located with a resolution of seven amino acids; corresponding to amino acids 146-152 of that protein. Then, a complete replacement set of peptides in which all 20 amino acids were substituted in turn at every position within the epitope was synthesized, and the particular amino acids conferring specificity for the reaction with antibody were determined. It was found that the leucine residues at positions 148 and 151 were essential for reaction with antisera raised against intact virus. A lesser contribution was derived from the glutamine and alanine residues at positions 149 and 152, respectively. Aside from the practical significance for locating and examining epitopes at high resolution, these findings may lead to better understanding of the basis of antigen-antibody interaction and antibody specificity.

Recombinant DNA technology now makes possible by deduction from the determined nucleotide sequences reliable amino acid sequences of biologically important proteins. However, methods for identifying the loci in a protein that constitute the antigenic and immunogenic epitopes are few and time consuming and form the bottleneck to further rapid progress. Immunogenic epitopes are defined as those parts of a protein that elicit the antibody response when the whole protein is the immunogen. These immunogenic epitopes are believed to be confined to a few loci on the molecule (1-3). On the other hand, a region of a protein molecule to which an antibody can bind is defined as an antigenic epitope. Antisera prepared against chemically synthesized peptides corresponding to short linear tracts of the total polypeptide sequence have been shown to react well with the native protein (4-9). However, interactions were also found to occur even when the site of interaction did not correspond to an immunogenic epitope of the native protein. This has been interpreted to mean that the number of immunogenic epitopes of a protein is less than the number of antigenic epitopes (4). Conversely, since antibodies produced against the native protein are, by definition, directed to the immunogenic epitopes, it follows that peptides reacting with these antibodies must contain elements of the epitopes. From a study of the few proteins for which the determinants have been accurately mapped, it is postulated that a determinant may consist of a single element (continuous) or of more than one element brought together from linearly distant regions of the poly-

peptide chain by the folding of that chain as it exists in the The publication costs of this article were defrayed in part by page charge

payment. This article must therefore be hereby marked "advertisement" in accordance with 18 U.S.C. \$1734 solely to indicate this fact.

The grant of the control of the cont native state (discontinuous) (10). Systematic mapping of all the detectable reactive elements of a protein by the chemical synthesis of overlapping segments has until now been se verely limited by the scale of the synthetic and testing capa bility required (10, 11) Smith and co-workers (12, 13) cicumvented the decoupling and purification steps by combin ing solid-phase peptide synthesis and solid-phase radioir munoassay using the same solid support:

We describe here the concurrent synthesis of all 208 pos ble overlapping hexapeptides covering the total 213-amin acid sequence of the immunologically important coat protein (VP1) of foot-and-mouth disease virus (FMDV), type 0 (Fig. 1). The peptides, still attached to the support used fo their synthesis, were tested for antigenicity by an ELISA using a variety of antisera. After identification of a hexaper tide reactive with antibody raised against the intact virus; all 120 hexapeptides representing the complete single point am. no acid replacement set were synthesized and tested for re tention of antigenicity. By this method a whole virus epitopwas examined at a resolution of a single amino acid.

#### MATERIALS AND METHODS

Synthesis of Peptides. Polyethylene rods (diameter, 4 mm length, 40 mm) immersed in a 6% (vol/vol) aqueous solution of acrylic acid were γ irradiated at a dose of 1,000,000 rads in rad = 0.01 gray) (15). Rods so prepared were assembled into a polyethylene holder with the format and spacing of a microtiter plate. Subsequent reactions at the tips of the rods were carried out in a Teflon tray with a matrix of wells to match the rod spacing. Conventional methods of solid-phase peptide chemistry (16, 17) were used to couple  $N^{\alpha}$ -t-buty! oxycarbonyl-L-lysine methyl ester to the polyethylene/poly acrylic acid via the Nº-amino group of the side chain Carboxy substitution of the support was determined by treating NH2-lysine(OMe)-polyethylene/polyacrylic acid with <sup>14</sup>C-labeled butyric acid and was found to be 0.15-0.2 nmol/mm<sup>2</sup>. Removal of the t-butyloxycarbonyl group was followed by the coupling of t-butyloxycarbonyl-L-alanine to complete a peptide-like spacer. Successive amino acids were added as dictated by the sequence to be synthesized. At the completion of the final coupling reaction, and after removal of the t-butyloxycarbonyl protecting group, the terminal amino group was acetylated with acetic anhydride in dimeth ylformamide/triethylamine. All N,N-dicyclohexylcarbodi, mide-mediated coupling reactions were carried out in di methylformamide in the presence of N-hydroxybenzotriazole. The following side-chain protecting groups were used O-benzyl for threonine, serine, aspartic acid, glutamic acid; and tyrosine; carbobenzoxy for lysine; tosyl for arginine; methylbenzyl for cysteine; and 1-benzyloxycarbonylamido-2,2,2-trifluoroethyl for histidine. Side-chain-protecting groups were removed by treatment with borontris(trifluors

Abbreviations: FMDV, foot-and-mouth disease virus; Pi/NoCli phosphate-buffered saline.

-T-T-S-A-G-E-S-A-D-P-( 201 amino acids )-T-L-COOH

# Best Available Copy

213

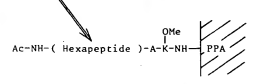


FIG. 1. The 213-amino acid sequence of VP1 (FMDV, type O<sub>1</sub>) stranslated by Kurz et al. (14) was subdivided into hexapeptide finits, and each was synthesized on a separate polyethylene support mithe orientation, and with a dipeptide spacer, as shown. Peptides are numbered according to the position of the NH-terminal amino acid within the VP1 sequence. PPA, polyethylene/polyacrylic acid.

cetate) in trifluoroacetic acid for 90 min at room temperature (18). After hydrolysis with HCl/propionic acid, sequences included in the synthesis as controls were analyzed confirm that, although coupling at each stage had occurred, it was incomplete for several of the amino acids, notably, arginine. Before testing by ELISA, support-coupled ceptides were washed several times with phosphate-buff-ted saline (P<sub>i</sub>/NaCl).

Antisera. Antisera against the intact virus particle were prepared by immunizing rabbits with 50  $\mu$ g of inactivated, density gradient-purified virus in complete Freund's adjuvant: The animals were bled 3-4 weeks after the single inoculation. Anti-virus-subunit serum was prepared by inocuating rabbits three times, 3-4 weeks apart, with 10  $\mu$ g of cid-disrupted purified virus, initially in complete Freund's and subsequently in incomplete Freund's adjuvant. The polypeptide VP1 was separated from the mixture of proteins ibtained from urea-disrupted purified virus by isoelectric fousing (19). It was eluted from the gel with 8 M urea and dalyzed against Pi/NaCl; and antiserum was raised in rabbits as described for the virus subunit. Antiserum for scan 3 see Fig., 2) was that used for scan 2; after absorption with purified virus (1500 µg of whole virus was incubated with 1 miof serum for 72 hr, at 4°C), and all virus-bound antibodies vereiremoved by centrifugation and an area of the area of the

EEISA. Support-coupled peptides were precoated with horse serum/10% ovalbumin/1% Tween 80 in Pi/NaCl or 1 hr. at 37°C to block nonspecific absorption of antibod-Overnight, incubation, at, 4°C, in, a 1:40 dilution, of antisein the preincubation mixture was followed by three Tashes in 0.05% Tween 80/P/NaCl Reaction for 1-hr-at-C with a:1:50,000 dilution of goat anti-rabbit IgG coupled Oborseradish peroxidase in the preincubation mixture was gain followed by extensive washing with P/NaCl/Tween to move excess conjugate. The presence of antibody was deested by reaction for 45 min with a freshly prepared develing solution (40 mg of o-phenylenediamine and 20  $\mu$ l of rogen peroxide in 100 ml of phosphate buffer, pH 5.0), indithe color produced was read in a Titertek Multiskan Laboratories, Melbourne, Australia) at 420 nm. Prior Offetesting, bound antibody was removed from the peptides washing peptides three times at 37°C in 8 M urea/0.1% 2nercaptoethanol/0.1% sodium dodecyl sulfate and then sevtimes with P<sub>i</sub>/NaCl.

#### RESULTS

dentification of a Virus Particle-Associated Immunogenic

sequence of the VP1 protein of FMDV type O1 were synthesized in duplicate. The amino acid sequence had been deduced from the nucleotide sequence of the VP1 gene (14). The results obtained for all the synthesized hexapeptides when tested by ELISA with six different antisera are shown in Fig. 2. Antisera used in the test were as follows: two different anti-(intact virus, type O<sub>1</sub>), a virus-absorbed anti-(intact virus, type  $O_1$ ), an anti-(virus subunit, type  $O_1$ ), an anti-(isolated virus protein VP1, type O<sub>1</sub>), and, as a control, an anti-(intact virus, type C<sub>3</sub>). The two anti-intact virus sera tested, scans 1 and 2, show the extremes in the reactivity patterns found. Large quantitative differences in the individual animal responses to an identical antigen preparation have been reported before, but these scans highlight the variability possible in the antibody composition between sera. Examination of scans 1, 2, and 3 shows that antibodies reactive with hexapeptide numbers 146 and 147 are present in antiintact particle sera (scans 1 and 2) but completely absent after absorption of the sera with purified virus (scan 3). Presumably, scan 3 registers those antibodies raised against epitopes expressed in denatured virions that are not present on the surface of the intact virion. Activities to hexapeptides 146 and 147 were not observed in the anti-subunit serum (scan 4) and were only weakly present in the anti-VP1 serum (scan 5). That some activity was found in the anti-VP1 serum

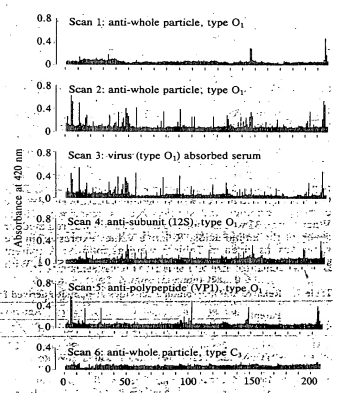


FIG. 2: Antigenic profiles (scans). Results are shown as vertical lines proportional to the extinction obtained in the antibody-binding ELISA test, plotted above the number giving the location within the VP1 sequence of the NH<sub>2</sub>-terminal amino acid of each peptide. Antisera used to produce the scans shown were as follows: 1 and 2, two different anti-whole virus particle, type  $O_1$ ; 3, anti-whole virus particle (as used in 2) after absorption with purified intact virus; 4, anti-virus subunit, type  $O_1$ ; 5, anti-VP1, type  $O_1$ ; 6, anti-whole virus particle, type  $C_3$ . It should be noted that, because the sequence of VP1 contains 20 alanine residues, 20 of the peptides synthesized match for seven amino acids. However, the frequency of reactive peptides from this group was not significantly different from the overall frequency (0.2 compared with 0.16) and therefore not considered further.

とは、それる はまた、いなななって、とはから、まなかってはん

possibly accounts for the immunizing capacity, albeit weak, of the isolated protein (20). It should be noted however that another anti-VP1 serum tested, while retaining a strong activity at position 148, showed no activity at positions 146 and 147. Comparison of scan 3 with scan 2 (absorbed compared with nonabsorbed) shows that, in addition to the loss of activity to peptides 146 and 147, some reduction in activity to peptides 5, 6, and 206 also occurred. Of these, activity to 5 and 6 was not found in all the anti-intact virus sera tested, but activity to 206 was invariably present. From this we conclude that of the peptides found to be reactive, the pair at 146 (G-D-L-Q-V-L) and 147 (D-L-Q-V-L-A) [in this paper, amino acids are identified by the single-letter code (21)] constitute or are part of the principal immunogenic epitope, with the element at 206 (V-A-P-V-K-P) contributing to a lesser epitope. This is consistent with the observations of others (5, 22) Scan 6 shows the absence of reactivity in an antiserum produced against a different serotype of the virus

Extending the Resolution of the Epitope at Peptides 146/147 to a Single Amino Acid From the preceding data, we were unable to distinguish between two possibilities: (i) the epitope is contained in the five amino acids common to peptides 146 and 147—i.e., D-L-Q-V-L—or (ii) the epitope is represented by the "sum", of the two hexapeptides—i.e., G-D-L-Q-V-L-A. To extend the resolution all 120 possible hexapeptides differing from peptide 146 (G-D-L-Q-V-L) by only a single amino acid were synthesized. Each of the other 19 common amino acids was substituted in each of the six amino acid positions within the peptide. Positions at which all or at least the majority of substitutions result in a loss of antibody-binding activity indicate those residues that are important for the specificity and binding to antibody. The ELISA activity obtained for each of the 120 peptides when serum 48 (anti-intact virus particle) was used in the test are shown in Fig. 3. The relative activities (with respect to the parent sequence) determined for each peptide for two different antiintact virus sera, nos. 31 and 48, are given in Table 1. To determine the contribution of the alanine residue (carboxyl terminus of peptide 147) toward reactivity and/or specificity, a further 20 peptides were synthesized. Each of these peptides consisted of the complete sequence of 146 (G-D-L-Q-V-L) with one of the 19 possible amino acids added to the carboxyl terminus and synthesized as described before. When serum 31 was used in the test, activity was retained for seven of the amino acids. Relative values expressed in the same way as given in Table 1 were as follows: A (parent amino acid), 99; D, 55; E, 36; G, 45; N, 95; Q, 98; S, 44. With

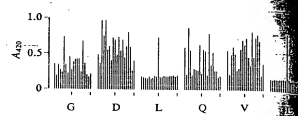


FIG. 3. Antibody-binding activity. The result for each period shown as a vertical line proportional to the ELISA extinction tained. Every group of 20 lines corresponds to the complete represent set for one of the six amino acid positions in the hexaper G-D-L-Q-V-L. Within each group of 20 lines, the left-hand-line responds to the substitution of the original residue by alaning and the successive lines are then in alphabetic order according to single letter code for the amino acids of the substitution of the original residue by alaning and the successive lines are then in alphabetic order according to single letter code for the amino acids of the substitution of the original residue by alaning and the successive lines are then in alphabetic order according to the substitution of the original residue by alaning acids of the substitution of the original residue by alaning acids

serum 48, activity was retained for four amino acids: A mentiamino acid); 94; G, 30; S), 47; T, 39 and a management and a mentiamino acid); 94; G, 30; S), 47; T, 39 and a management and a manag

#### 南京、中国中国的一部中国DISCUSSION 中央的自由

Interpretation of Data: In choosing to adopt the process for peptide synthesis as described; we made several assumitions:

1. To detect antibodies, the quantity of peptide of a fined sequence need only be in the pmol range (5). Assume a worst-case overall yield of 1% for an eight-step synthety (two linking and six sequence amino acids), an initial level 1 nmol of reactive group per support would satisfy the abscendition.

2. High purity for the peptide used in the detection of bodies is not a necessary condition. The majority of service cal tests rely on the specificity of antibodies to detect a antigen in the presence of large amounts of irrelevant potein.

3. Except for cases in which either all or none of the tides react, a large number of the peptides would effective act as negative controls in the test. With adjacent peptides sharing a common sequence of five amino acids, the observation of peaks above a generally uniform background lewould indicate a valid test.

4. Many of the antibodies elicited by immunization wan intact virus result from presentation of epitopes in fully partially denatured form. Such antibodies may bind to such thetic peptides in vitro but not to the virion itself. They therefore assumed to be less relevant to virus neutralization.

Table 1. Relative antibody-binding activities of peptides derived from the parent sequence G-D-L-Q-V-L

Serum	Parent residue	Activity when substituted with animo acid															.74			
		A	С	D	Е	F	G	Н	I	K	L	M	N	P	Q	R	S	Т	V	Ŵ
31	G	29					90	14	27	12		32	34	41	29		50			. 70
	D	22	21	143	95	110	28	65		12	65	15	58	10	69		38	62		-1
	L						•				79									1
	Q			64	14								13		80					
	v	62		33	52				26		29	59			45		49	43	89	
	L										119		***	•• .						
48	G	11					88	10		32		18	24	25	26		77	14		
	D	37	12	136	92	137	52	62	21	87	81	37	89	49	80	29	63	104	60	
	L										88									
	Q	60		117	52					68		53	49		102	10	45			
	V	52		40	63	42			56	82	68	88	34		106	33	91	98	81	-
	L										105									

Antibody-binding activities are shown for all peptides that gave an extinction significantly above background. Values for each peptide expressed as a percentage of the mean activity of the six parent sequences synthesized as a part of each replacement set. Values given bolding type correspond to those obtained for the parent sequence. No activity was detected when the antiserum used was prepared against heterologous FMDV type.

than are antibodies that bind to virions as well as to peptides. The extinction obtained in an ELISA for a given peptide depends on the concentration of the antibody population with the correct specificity for reaction. It is essentially independent of the peptide density expressed as reacting groups mm<sup>2</sup> of support (unpublished data). The difference in the extinction obtained with peptides synthesized with densities varying over two orders of magnitude is similar to the 10-20% variation observed between replicate synthesis (unpubished data). The extinction may also be expected to depend greatly on the affinity between peptide and reacting antibody, but this remains to be verified, although the overnight reaction would tend to minimize differences. Antigenic profiles of the FMDV VP1 (Fig. 2) were interpreted to define an antigenic peptide as one giving an ELISA extinction significantly above the background level of the test. On the other hand, in the testing of replacement nets (Fig. 3), the concenfration of the reactive antibody population is constant and effectively of one specificity. Therefore, the extinctions observed are interpreted to reflect the mean affinity of the reacting antibody population for the peptide.

Immunogenicity of FMDV Virus Protein (VP1). Scan 5 of Fig. 2 identifies six immunogenic regions defined in terms of epitopes on the isolated protein eliciting antibodies capable of binding to the corresponding synthetic peptide. Scan 4 shows that, for the same protein as a part of the virus subunit, additional regions (principally 50-70 and 191-197) are immunogenic. Scan 2 shows that, during the course of the mmune response to whole virus, most of the protein can be immunogenic. In contrast, scan 1 shows a response to only a very limited number of epitopes. What has become clear from these and other results (unpublished) is that different animals do not necessarily respond to all of the epitopes on a given antigen. In addition, the immunogenic response of an individual animal will be complicated if the antigen is readily broken down as is known to happen to FMDV (23, 24). The animal is exposed not only to the intact virus but also to subunits and possibly even to the isolated viral proteins. Each of these different states could present different epitopes to the immune system. Epitopes can be identified with a particular state of the antigen by testing the peptides with antisera specific to that state.

An Immunogenic Epitope at High Resolution. Antibodies aised against a particular immunogenic epitope will have a combining site (paratope) complementary to the structure of that epitope. An antibody population directed to the same epitope (allowing for variation in the expression of antibodies by the immune reponse) will have common features in the combining sites essential for binding to that epitope. A peptide that, in one of its many conformations in thermal equilibrium in vitro, has a structure sufficiently similar to the form of the epitope against which antibody was raised in vivo will bind to the antibody. Modification of a reacting peptide by amino acid substitution will define the limits for interaction with antibody. By so "mapping" the antibody-combining site, it is possible to infer properties of the antigen to which this antibody population is complementary. Using polyclonal antisera, it was not expected that a rigorous requirement for particular amino acids in particular positions would be observed. It is clear that, whatever the diversity of the antibodies involved in the interaction, the requirement for a given amino acid in certain positions is absolute for most or all of the antibodies present. It is also clear that the specificity range found for the two different antisera is remarkably similar, differing mainly in the hierarchy of preference for amino acids at the nonessential position. As judged from the limitation to replacements at some position within the sequence G-D-L-Q-V-L-A, the whole-virus epitope may be considered to be X-X-L-Q-X-L-A, where X is nonessential, letters in boldface type indicate an absolute requirement, and letters in lightface type indicate a contributing amino acid.

4001

These findings suggest a different interpretation of the characteristics of epitopes. The antigenic specificity of the epitope represented by amino acids 146-152 within the VP1 protein of FMDV is largely dependent on the leucine residues at positions 148 and 151. These are hydrophobic residues and would not normally be expected to protrude from the protein surface. This suggests the possibility that the immune system responds to a local protein conformation that is different from that expected to represent the global energy. minimum. The energy for antigen-antibody binding may be derived from the positive entropy term associated with the transfer of hydrophobic residues from a hydrophilic (aqueous) environment to within the antibody-combining site.

Scope of the Described Approach to Epitope Mapping. Although our results have been presented for a single protein only, the agreement with results of others in locating a viral epitope within the region encompassing amino acids 141-160 of VP1 is excellent (5, 22). The further resolution obtained by Rowlands et al. (25) from the comparison of the sequences of the VP1s of three antigenic variants of a single virus type  $(A_{12})$  showed that amino acid substitution at positions 148 and/or 153 would affect the ability to react with specific antibody. This result is in good agreement with our results for subtype O<sub>1</sub>, where positions 148 and 151 were critical to the immunogenicity of the epitope. We expect that the systematic approach as outlined, when applied to a broader spectrum of proteins, will contribute greatly to our understanding of the nature of epitopes and their interaction with the immune system.

We thank Mr. Jan Briaire for his enthusiastic and skilled technical assistance with the synthesis of the peptides, Mr. Jan Meyer for assistance with the ELISA, and Dr. Dick Voskamp of the Technical University, Delft, for valuable advice on aspects of the peptide chemistry. This work is the result of a collaborative project between the Commonwealth Serum laboratories and the Central Veterinary Institute and was conducted at the latter institute:

- Crumpton, M. J. (1974) in The Antigens; ed. Sela, M. (Aca-رين برورين demic, New York); الإولى demic, New York) و المراجعة ا
- Benjamini, E., Michaeli, D. & Young, J. D. (1972) Curr. Top. Microbiol. Immunol. 58, 85-134.
- 3. Atassi, M. Z. (1975) Immunochemistry 12, 423-438.
  4. Green, N., Alexander, H., Olson, O., Alexander, S., Shinnick, T. M., Sutcliffe, J. G. & Lerner, R. A. (1982) Cell 28, 477-487.
- 5. Bittle, J. L., Houghten, R. A., Alexander, H., Shinnick, ToMar Sutcliffe, J. G., Lerner, R. A., Rowlands, D. J. &
- 6. Dreesman, G. R., Sanchez, Y., Ionescu-Matiu, I., Sparrow, J. T., Six, H. R., Peterson, D. L., Hollinger, F. B. & Melnick, J. L. (1982) Nature (London) 295, 158-160.
- 7 Prince, A. M., Ikram, H. & Hopp, T. P. (1982) Proc. Natl. TAcad: Sci.: USA: 79; 579=582.7 134
- .. 821, Lerner, R.: A., Green, N., Alexander, H., Liu, F., Sutcliffe, J.G. & Shinnick, T. M. (1981) Proc. Natl. Acad. Sci. USA 78, 3403-3407.
- Neurath, A. R., Kent, S. B. H. & Strick, N. (1982) Proc. Natl. Acad. Sci. USA 79, 7871-7875.
- 10. Atassi, M. Z. (1978) Immunochemistry 15, 909-936:
- 11. Kazim, A. L. & Atassi, M. Z. (1980) Biochem. J. 191, 261-
- Smith, J. A., Hurrell, J. G. R. & Leach, S. J. (1977) Immunochemistry 14, 565-568.
- Hurrell, J. G. R., Smith, J. A. & Leach, S. J. (1978) Immunochemistry 15, 297-302. Kurz, C., Forss, S., Kupper, H., Strohmaier, K. & Schaller,
- H. (1981) Nucleic Acids Res. 9, 1919-1931. Muller-Schulte, D. & Horster, F. A. (1982) Polym. Bull. 7, 77-
- Erickson; B. W. & Merrifield, R. B. (1976) in The Proteins, eds. Neurath, H. & Hill, R. L. (Academic, New York), Vol. 2, pp. 255-527.5

Best Available Copy

Biochemistry: Geysen et al.

Proc. Natl. Acad. Sci. USA 81 (1984)

4002

- 17. Meienhofer, J. (1973) in Hormonal Proteins and Peptides, ed. Li, C. H. (Academic, New York), Vol. 2, pp. 45-267.
- Pless, J. & Bauer, W.: (1973) Angew. Chem. 85, 142.
- Barteling, S. J., Wagenaar, F. & Gielkens, A. L. J. (1982) J. Gen. Virol. 62, 357-361.
- Kleid, D. G., Yansura, D., Small, B., Dowbenko, D., Moore, D. M., Grubman, M. J., McKercher, P. D., Morgan, D. O., Robertson, B. H. & Bachrach, H. L. (1981) Science 214,

हुँ । १९४५, १९०० - १९४४ - १९०० - १९०७ मध्ये १९०७ व्यक्तिक व्यक्ति । १९४५ - १९०० व्यक्ति । १९४४ - १९०० - १९०० -१९४४ - १९४५ - १९४४ - १९४४ - १९४४ - १९४४ - १९४४ - १९४४ - १९४४ - १९४४ - १९४४ - १९४४ - १९४४ - १९४४ - १९४४ - १९४४ भिन्न के <mark>प्रस्तानेत उ</mark>न्होंने, के ज़ार में हैं स्ट्रिक्ट **में के** सामक्षेत्रकार हो। अपूर्व Tarabesturi entroduction of the continuous of the feet of the second of the second sec mandage in a construction of the dealers of the construction of the dealers of th reference to tradical to see that arrest to a little field in the filling Committee of the state of the s Mi & fire ्राच्या है जो का कार्य है है कि कार्य है की स्थाप े राज्य । १ । १६ । १४८ ह्यु स्क्रुब्रे र या गाम देख्या क्रांके स्वय

21. IUPAC-IUB Commission on Biochemical Nomencia (1968) Eur. J. Biochem. 5, 151-153.

-Pfaff, E., Mussgay, M., Bohm, H. O., Schulz, G. E Schaller, H. (1982) EMBO J. 1, 869-874.

Brown, F. (1981) Trends Biochem. Sci. 6, 325-327.

Meloen, R. H. & Briaire, J. (1980) J. Gen. Virol. 55, 107-11. Rowlands, D. J., Clarke, B. E., Carroll, A. R., Brown, Nicholson, B. H., Bittle, J. L., Houghten, R. A. & Leme R. A. (1983) Nature (London) 306, 694-697.

in and the strong and the second of the seco

lan tilland. I nomin oli yustidh svoquotsaktin **no**n

सर्वे अपना शामानामाना प्रतिकृति होता । द्वारा स्थाप स्थापना स्थापना स्थापना ।

ed are interpreted to return common titlery more interpret

The artification of the company of the confidence of the confidenc

Calculations are in the second to be a second consequent with a produced